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- Block polyesters having branched hydrophilic capping groups useful as soil release agents in detergent compositions.
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Description

Technical Field

The present application relates to block polyesters and like compounds having branched hydrophilic capping groups which are useful as soil release agents in laundry detergent compositions.

In addition to cleaning performance, laundry detergent compositions desirably have other benefits. One is the ability to confer soil release properties to fabrics woven from polyester fibers. These fabrics are mostly co-polymers of ethylene glycol and terepithalic acid, and are sold under a number of tradenames, as e.g. Dacron, Fortrel, Kodel and Blue C Polyester. The hydrophobic character of polyester fabrics makes their laundering difficult, particularly as regards oily soil and oily stains. The oily soil or stain preferentially "wests" the fabric. As a result, the oily soil or stain is difficult to remove in an anguesus laundering corceasts.

High molecular weight (e.g., 40.000 to 50.000 M.W.) polyesters containing random ethylene terephthalate/polyethylene glycol (PEG) terephthalate units have been used as soil release compounds in laundry 15 detergent compositions. See U.S. Patent 3,962,152 to Nicol et al., issued June 8, 1976. During the laundering operation, these soil release polyesters adsorb onto the surface of fabrics immersed in the wash solution. The adsorbed polyester then forms a hydrophilic film which remains on the fabric after it is removed from the wash solution and dried. This film can be renewed by subsequent washing of the fabric with a delergent composition containing the soil release polyesters.

These ethylene terophthalate/PEG terephthalate polyesters are not very water-soluble. It is believed that they form a suspension in the wash solution which does not adsorb efficiently onto the fabrics. As a result, the level of soil release polyester in the detergent composition has to be increased if benefits are to be obtained after several wash cycles. Because of this poor water-solubility, these polyesters are formulated as suspensions in laundry detergent compositions, rather than as isotropic liquids. In certain detergent formulations, these polyesters can also diminish clay soil cleaning performance.

Background Art

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A. Polyester anti-static agents formed from dimethyl terephthalate, ethylene glycol and methoxy PEGs.

U.S. Patent 3,416,962 to McIntrye et al., issued December 17, 1988, discloses the treatment of shaped polyester articles with a water-insoluble crystallizable polymeric compound which can contain a water-solvatable polymeric group such as a polyoxyalkylene group having an average molecular weight of from 300-0000. Preferred polyoxyalkylene groups are the PEGs having an average molecular weight of from 1000-4000. Treatment of the shaped articles is carried out by applying an aqueous dispersion of the crystallizable polymeric compound in the presence of an anti-oxidant, followed by heating to a temperature above 90 °C to obtain a durable coating of the compound on the shaped article. Example 6 discloses one such crystallizable polymeric compound formed by the reaction of dimethyl terephthalate, ethylene glycol and an _O-methyl poly(oxyethylene.) glycol of average molecular weight 350. A 20% solution of this polysets in benzyl alcohol was used to impart anti-static properties to a polyseter fabric. Example 7 discloses a 20% aqueous solution of a similar polyseter used to impart anti-static properties to a polysets to the control of the control of

B. Polyester anti-static and soil release agents formed from dimethyl terephthalate, sodium dimethyl 5sulphoisophthalate, ethylene glycol and polyethylene glycol (PEG)

U.S. Patent 4,427,587 to Stockburger, filed February 15, 1983, issued January 24, 1984, discloses low molecular weight copolyesters (M.W. 2,000 to 10,000) which can be used in aqueous dispersions to impart soil release properties to polyester fibers. The copolyesters are formed by the reaction of ethylene glycol, a 50 PEG having an average molecular weight of 200 to 1000, an aromatic dicarboxylic acid (e.g. dimethyl terephthelate), and a suffonated aromatic dicarboxylic acid (e.g. dimethyl 5-sufficisphthalate). The PEG can be replaced in part with monoalkylethers of PEG such as the methyl, ethyl and butyl ethers. A dispersion or solution of the copolyester is applied to the textile material and then heat set at elevated temperatures (90* to 150* C1 to impart durable soil release properties.

C. Monomeric polyesters of PEG and terephthalic acid useful as soil release agents

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U.S. Patent 4,349,688 to Sandler, issued September 14, 1982, discloses polyoxyalkylene ester soil release agents, in particular monomeric polyesters of PEG and terephthalic acid having the formula:

where n can range from 6-23 and X is either methyl or H. Example IV discloses the preparation of one such PEG(terephthalate polyester formed from terephthaloyl chloride and Carbowax 400 (n = 9, X = H). Durable soil resistancy and water wicking properties are imparted by wetting the fabric with a composition to containing the polyoxyalkylene ester, drying the wetted fabric, and then curing the dried fabric at a temperature of from 190 '-200 ' C for about 45-90 seconds.

D. Ethylene terephthalate/PEG terephthalate soil release polyesters for fabric treating solutions.

20 U.S. Patent 3,959,230 to Hays, Issued May 25, 1976, discloses polyester soil release agents containing random ethylene terophthalate/PEG terephthalate units in a mole ratio of from about 25.75 to about 35.55. These soil release polyesters have a molecular weight of from about 25.000 to about 55.000, (predestly from about 40,000 to about 55,000) and are used in dilute, aqueous solutions, preferably with an emulsifying agent present. Fabrics are immersed in this solution so that the soil release polyester adsorbs onto the 25 fabric surface. The polyester forms a hydrophilic film which remains on the fibers after the fabric is removed from the solution and dried. See also U.S. Patent 3,893,929 to Basadur, issued July 8, 1975 (compositions for imparting soil release finish containing a polyester having an average molecular weight of 3000-5000 formed from terephthalic acid, PEG and ethylene glycol); U.S. Patent 3,712,873 to Zenk, issued Junuary 23, 1973 (textile treating composition comprising fatty alcohol polyethoxylates; quaternary ammonium compounds; a polyester having average molecular weight of 3000-5000 formed from terephthalic acid. PEG and ethylene glycol; and starch).

E. Ethylene terephthalate/PEG terephthalate soil release agents used in detergent compositions.

U.S. Patent 3,982,152 to Nicol et al., issued June 8, 1976, discloses detergent compositions containing detergent surfactants and the ethylene terophthalate/PEG terophthalates soil release polyesters disclosed in the Hays patent. See also U.S. Patent 4,116,885 to Derstadt et al., issued September 28, 1978 (detergent compositions containing certain compatible anionic detergent surfactants and ethylene terephthalates/PEG terophthalate soil release polyesters); U.S. Patent 4,132,880 to Nicol, issued January 2, 1979 (detergent ownositions containing detergent surfactants; a composition which disassociates to yield quaternary ammonium cations; and an ethylene terephthalate/PEG terophthalate/PEG tierophthalate/PEG tierophthalate

F. Soil release and antistatic polyurethanes useful in detergent compositions which contain polyester blocks having sulfoisophthalate units.

U.S. Patent 4,201,824 to Violland et al., issued May 6, 1980, discloses hydrophilic polyurethanes having soil release and antistatic properties useful in detergent compositions. These polyurethanes are formed from the reaction product of a base polyester with an isocyanate prepolymer (reaction product of diisocyanate and macrodiol). Example VI discloses a base polyester formed from dimethyl terephthalate, 50 dimethyl sulfosophthalate, ethylene glycol and PEG (molecular weight 300) which is reacted with a prepolymer formed from a PEG (molecular weight 1,300) and toluene diisocyanate.

DISCLOSURE OF THE INVENTION

The present invention relates to compounds of formula:

 $X_{\{(OCH_2CH(Y))_n(OR^5)_m\}}_{\{(A-R^1-A-R^2)_u(A-R^3-A-R^2)_v\}}$ -A-R⁴-A $_{\{(R^5O)_m(CH(Y)CH_2O)_n\}}_{\{(A-R^1-A-R^2)_u(A-R^3-A-R^2)_v\}}$ wherein the A moieties are essentially



moieties; the R¹ moieties are essentially 1,4-phenylene moieties; the R² moieties are essentially ethylene to moieties, or substituted ethylene moieties having C₁-C₁ alkyl or alkoxy substituents; the R² moieties are substituted C₂-C₁s hydrocarbylene moieties having at least one -SO₃M, -COOM, -OŢ(R²O)m(CH(Y)CH₂O)-₃1X or

-A{(R²-A-R¹-A)}_w{(R²O)_m(CH(Y)CH₂O)_m}X substituent or at least one molety -A{(R²-A-R¹-A)}_wR²-A crossitinked to another R² molety; the R² moleties, or a mixture thereof; each R² is C₁-C₄ allyleine, or the molety -R²-A-R², where R² is a C₁-C₁-C₂ allyleine, allyleinyleine, aryleine or alkaryleine molety; each M is H or a water-soluble cation; the Y substituents of each molety {(R²O)_m(CH(Y)CH₂O)_m} are the ether molety -CH₂(OCH₂O+D_m)_mOX or a mixture of this ether molety and H; each X is H, C-C₂ alkylor

wherein R' is G_1 - G_2 alkyl; m and n are numbers such that the moiety -(CH(Y)CH₂O)- comprises at least about 50% by weight of the moiety -{(R*O)_m(CH(Y)CH₂O)_m}, provided that when R° is the moiety -R² -AR°- G_1 , m is 1; each n is at least about G_2 is 0 or at least 1; u and v are numbers such that the sum of u + v is from about 3 to about 25 when w is 0; w is 0 or at least 1; and when w is at least 1, u, v and w are numbers such that the sum of u + v + w is from about 3 to about 25.

The present invention further relates to detergent compositions, especially for laundry use, which so comprise a soil release component having an effective amount of these compounds. These detergent compositions further comprise from about 1 to about 75% by weight of a nonionic, anionic, ampholytic, zwitterionic, or cationic detergent surfactant, or mixture thereof. In addition to these detergent surfactants, the detergent compositions can optionally comprise from 0 to about 60% by weight of detergent buildings.

The compounds of the present invention provide excellent soil release benefits to polyseter fabrics of during laundering, but without diminishing the clay soil cleaning performance of the detergent composition. These compounds can be used at lower levels in detergent compositions to provide soil release benefits at least equivalent to prior art high molecular weight ethylene terephthalate/PEG terephthalate polyesters. Some of the compounds of the present invention can also be formulated to provide isotropic liquid detergent compositions.

Cortain of the soil release compounds of the present invention provide additional through-the-wash static control benefits to laundered fabrics. The compounds of the present invention also provide cleaning benefits in terms of greasy/oily stain removal, as well as whiteness maintenance benefits. In addition, it is expected that the soil release compounds of the present invention will be more biodegradable than prior art ethylene terephthaltate/EG terephthaltate post release polyesters.

Soil Release Compounds

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The compounds of the present invention have the formula:

In this formula, the moiety {(A-R1-A-R2)_w(A-R3-A-R2)_w} A-R4-A-forms the oligomer or polymer backbone of

the compounds. It is believed that the structure of the backbone is what is important to the actsorption of the compounds on polyester fabrics during laundering. Groups $X\{(CCh_ECH(Y))_m(DR^2)_m\}$ and $\{(R^SO)_m(CH(Y)-CH_2Oh_g)_m\}$ X are generally connected at the ends of the oligometripolymer backbone. It is believed that soil release properties of the compounds (when absorbed on the fabric) are due to these hydrophilic end 5 groups.

The linking A moieties are essentially



20 moieties, i.e. the compounds of the present invention are polyesters. As used herein, the term "the A moieties are essentially



mojeties" refers to compounds where the A mojeties consist entirely of mojeties

or are partially substituted with linking moieties such as

or

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an

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(amide), and

or

25 (urethane). The degree of partial substitution with these other linking moieties should be such that the soil release properties are not adversely affected to any great extent. Preferably, linking moieties A consist entirely of (i.e., comprise 100%) moieties

The R1 moieties are essentially 1,4-phenylene moieties. As used herein, the term "the R1 moieties are essentially 1,4-phenylene moieties" refers to compounds where the R1 moieties consist entirely of 1,4-55 phenylene moieties, or are partially substituted with other arylene or alkarylene moieties, alkylene moieties, alkenylene moieties, or mixtures thereof. Arylene and alkarylene moieties which can be partially substituted for 1,4-phenylene include 1,3-phenylene, 1,2-phenylene, 1,8-naphthylene, 1,4-naphthylene, 2,2-biphenylene, 4,4'-biphenylene and mixtures thereof. Alkylene and alkenylene moieties which can be partially substituted include ethylene, 1,2-propylene, 1,4-butylene, 1,5-pentylene, 1,6-hexamethylene, 1,7-heptamethylene, 1,8octamethylene, 1,4-cyclohexylene, and mixtures thereof.

For the R1 mojeties, the degree of partial substitution with mojeties other than 1.4-phenylene should be such that the soil release properties of the compound are not adversely affected to any great extent. Generally, the degree of partial substitution which can be tolerated will depend upon the backbone length of the compound, i.e., longer backbones can have greater partial substitution for 1,4-phenylene moieties. Usually, compounds where the R1 comprise from about 50 to 100% 1,4-phenylene moieties (from 0 to about 50% moieties other than 1,4-phenylene) have adequate soil release activity. For example, polyesters made according to the present invention with a 40:60 mole ratio of isophthalic (1,3-phenylene) to terephthalic (1,4-phenylene) acid have adequate soil release activity. However, because most polyesters used in fiber making comprise ethylene terephthalate units, it is usually desirable to minimize the degree of partial substitution with moieties other than 1.4-phenylene for best soil release activity. Preferably, the R1 moieties consist entirely of (i.e., comprise 100%) 1,4-phenylene moieties, i.e. each R1 moiety is 1,4phenylene.

The R2 moieties are essentially ethylene moieties, or substituted ethylene moieties having C1-C4 alkyl or alkoxy substitutents. As used herein, the term "the R2 moieties are essentially ethylene moieties, or substituted ethylene moieties having C1-C4 alkyl or alkoxy substituents" refers to compounds of the present invention where the R2 moieties consist entirely of ethylene, or substituted ethylene moieties, or are partially substituted with other compatible moieties. Examples of these other moieties include linear C3-C6 alkylene 20 moieties such as 1.3-propylene, 1.4-butylene, 1.5-pentylene or 1.6-hexamethylene, 1.2-cycloalkylene moieties such as 1,2-cyclohexylene, 1,4-cycloalkylene moieties such as 1,4-cyclohexylene and 1,4dimethylenecyclohexylene, polyoxyalkylated 1,2-hydroxyalkylenes such as

and oxyalkylene moieties such as -CH2CH2OCH2CH2OCH2CH2- or -CH2CH2OCH2CH2-.

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For the R2 moieties, the degree of partial substitution with these other moieties should be such that the soil release properties of the compounds are not adversely affected to any great extent. Generally, the degree of partial substitution which can be tolerated will depend upon the backbone length of the compound, i.e., longer backbones can have greater partial substitution. Usually, compounds where the R2 comprise from about 20 to 100% ethylene, or substituted ethylene moieties (from 0 to about 80% other 35 compatible moieties) have adequate soil release activity. For example, polyesters made according to the present invention with a 75:25 mole ratio of diethylene glycol (-CH2CH2OCH2CH2-) to ethylene glycol (ethylene) have adequate soil release activity. However, it is desirable to minimize such partial substitution, especially with oxyalkylene moieties, for best soil release activity. (During the making of polyesters according to the present invention, small amounts of these oxyalkylene moieties (as dialkylene glycols) are 40 typically formed from glycols in side reactions and are then incorporated into the polyester). Preferably, R² comprises from about 80 to 100% ethylene, or substituted ethylene moieties, and from 0 to about 20% other compatible moieties.

For the R2 moieties, suitable ethylene or substituted ethylene moieties include ethylene, 1,2-propylene. 1,2-butylene, 1,2-hexylene, 3-methoxy-1,2-propylene and mixtures thereof. Preferably, the R2 moieties are 45 essentially ethylene moieties, 1,2-propylene moieties or mixtures thereof. Inclusion of a greater percentage of ethylene moieties tends to improve the soil release activity of the compounds. Surprisingly, inclusion of a greater percentage of 1.2 propylene moieties tends to improve the water solubility of the compounds.

For the R3 moieties, suitable substituted C2-C18 hydrocarbylene moieties can include substituted C2-C12 alkylene, alkenylene, arylene, alkarylene and like moieties. The substituted alkylene or alkenylene moieties can be linear, branched, or cyclic. Also, the R3 moieties can be all the same (e.g. all substituted arylene) or a mixture (e.g. a mixture of substituted arylenes and substituted alkylenes). Preferred R3 mojeties are those which are substituted 1.3-phenylene mojeties.

The substituted R3 moieties preferably have only one - SO3M, -COOM, -Of (R5O)m(CH(Y)CH2O)n1X or -Af (R2-A-R4-A)1wf (R50)m(CH(Y)CH2O)m1 X substituent. M can be H or any compatible water-soluble cation. 55 Suitable water soluble cations include the water soluble alkali metals such as potassium (K+) and especially sodium (Na+), as well as ammonium (NH4+). Also suitable are substituted ammonium cations having the formula:

$$R^2 - \frac{R^1}{N^+} - R^4$$

where R¹ and R² are each a C·-C₂c hydrocarbyl group (e.g. alkyl, hydroxyalkyl) or together form a cyclic or heterocyclic ring of from 4 to 6 carbon atoms (e.g. piperidine, morpholine); R³ is a C·-C₂c hydrocarbyl group; and R⁴ is H (ammonium) or a C₁-C₂c hydrocarbyl group (quat amine). Typical substituted ammonium cationic groups are those where R⁴ is H (ammonium) or C₁-C₂ alkyl, especially methyl (quat amine); R¹ is C₁₀-C₁₂ alkyl, especially C₁₂-C₁₄ alkyl, and R² and R³ are each C₁-C₄ alkyl, especially

The R3 moieties having -Af (R2-A-R4A)1-

f (R² O)_m(CH/)(CH₂O)_m)X substituents provide branched backbone compounds. R³ moleties having -Af (R²-15 A-R²-A))_m-R²- notieties provide crosslinked backbone compounds. Indeed, syntheses used to make the branched backbone compounds typically provide at least some crosslinked backbone compounds.

The moleties (R⁶O) and -(CHY)CH₂O)- of the moleties {(R⁶O)_m(CHY)CH₂O)-} and {(COHY)CH₂O)-(CH⁶)_m} can be mixed together or preferably form blocks of -(R⁶O)- and -(CHY)CH₂O)-moleties. Preferably, the blocks of -(R⁶O)- moleties are located next to the backbone of the compound. When R⁶ is the 20 molety -R²-A-R⁶-, m is 1; also, the moiety -R²-A-R⁶ - is preferably located next to the backbone of the compound. For R⁶, the preferred C₂-C₂ alkylene is C₂-C₄ (propylene), when R⁶ is C₂-C₄ alkylene is preferably from 0 to about 5 and is most preferably 0. R⁶ is preferably methylene or 1,4-phenylene. The moiety -(CHY)CH₂O)- preferably comprises at least about 75% by weight of the moiety -{(R⁶O)_m(CHY)-CH₂O)₁, and most preferably 100% by weight (m is 0).

The Y substituents of each molety $\{(R^0O)_m(CH(Y)CH_0O)_n\}$ are the other molety $-CH_0(CCH_0)_pC-X$, or are, more typically, a mixture of this ether molety and H; p can range from 0 to 100, but is typically 0. When the Y substituents are a mixture, molety $-(CH(Y)CH_0O)_m$, can be represented by the following molety:

$$\begin{array}{c} -\mathbb{E}(\mathsf{CHCH}_2\mathsf{O})_{\mathsf{n}_1}(\mathsf{CH}_2\mathsf{CH}_2\mathsf{O})_{\mathsf{n}_2} \\ \\ \mathsf{CH}_2(\mathsf{OCH}_2\mathsf{CH}_2)_{\mathsf{D}}\mathsf{O}-\mathsf{X} \end{array}$$

wherein n_1 is at least 1 and the sum of $n_1 + n_2$ is the value for n. Typically, n_1 has an average value of from about 1 to about 10. The moieties

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can be mixed together, but typically form blocks of

and

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moieties. X can be H, C1-C4 alkyl or

wherein R' is C--C, alkyl. X is preferably methyl or ethyl, and most preferably methyl. The value for each n is at least about 6, but is preferably at least about 10. The value for each n usually ranges from about 12 to about 113. Typically, the value for each n is in the range of from about 12 to about 43.

The backbone moieties (A-R¹-A-R²) and -(A-R²-A-R²) and be mixed together or can form blocks of (A-R¹-A-R²) and (A-R³-A-R²) moieties. It has been found that the value of u + v needs to be at least about 3 in order for the compounds of the present invention to have significant soil release activity. The amaximum value for u + v is generally determined by the process by which the compound is made, but can range up to about 25, i.e. the compounds of the present invention are oligomers or low molecular weight, polymers. By comparison, polyesters used in fiber making typically have a much higher molecular weight, e.g. have from about 50 to about 250 ethylene terephthalate units. Typically, the sum of u + v ranges from about 30 to about 10 for the compounds of the present invention.

26 Generally, the larger the u + v value, the less soluble is the compound, especially when the R² moieties do not have the substituents -COOM or -SO_VM. Also, as the value for n increases, the value for u + v should be increased so that the compound will deposit better on the fabric during laundering. When the R² moieties have the substituent -A²(R²-AR²-A)_V-(R²O_M(CHY)O²H₂O_M)_AX (tranched backbone compounds), the value for w is typically at least 30 1 and is determined by the process by which the compound is made. For these branched and crosslinked backbone compounds the value for u + v + w is from about 30 to about 25.

Preferred compounds of the present invention are block polyesters having the formula:

$$\begin{array}{c} x - E \left(\text{OCH}_2\text{CH}_2 \right)_{n_2} \left(\text{OCH}_2\text{CH} \right)_{n_1} - \frac{0}{1} - \frac{0}{1} \left(\text{OC} - \text{R}^1 - \frac{0}{1} - \text{CO} - \text{R}^2 \right)_{\mathbf{u}} \left(- \text{OC} - \text{R}^3 - \frac{0}{1} - \frac{0}{1} - \frac{0}{1} \right)_{\mathbf{u}} \left(- \text{OC} - \frac{0}{1} - \frac$$

wherein the R¹ moieties are all 1.4-phenylene moieties; the R² moieties are essentially ethylene moieties, 1.2-propylene moieties or mixtures thereof; the R³ moieties are all potassium or preferrably sodium 5-sulfo-1.3-phenylene moieties or substituted 1.3-phenylene moieties having the substituent

$$\bigcap_{-CO-E-(R^2-OC-R^4-CO)-\frac{1}{W}-E(CHCH_2O)_{n_1}(CH_2CH_2O)_{n_2}-X \text{ at the } 5$$

position; the \mathbb{R}^4 moleties are \mathbb{R}^1 or \mathbb{R}^3 moleties, or mixtures thereof; each X is ethyl or preferably methyl; each n; is from 1 to about 5; the sum of each n; $+ n_2$ is from about 12 to about 43; when w is 0, $+ n_3$ to $+ n_3$ to + n

Particularly preferred block polyesters are those where v is 0, i.e. the linear block polyesters. For these most preferred linear block polyesters, u typically ranges from about 3 to about 8. The most water soluble of these linear block polyesters are those where u is from about 5 to about 5.

Method for Making Compounds

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The compounds of the present invention can be prepared by art-recognized methods. Although the following synthesis description is for the preferred block polyesters of the present invention, other versions can be prepared by appropriate variation.

The block polyesters of the present invention are typically formed from: (1) ethylene glycol, 12propylene glycol or a mixture thereof; (2) a capping group formed from the reaction of the appropha-15 glycidyl ether with a polyethylene glycol (PEG) capped at one end with a C₁-C₄ alkyl group; (3) a dicarboxylic acid (or its diester), and optionally (4) an alkall metal salt of a sulfonated aromatic dicarboxylic acid (or its diester), or if branched polyesters are desired, a polycarboxylic acid (or its esten). The respective amounts of these four components are selected to prepare polyesters having the desired properties in terms of solubility and soil release properties.

The capped PEG used to prepare the capping group is typically methyl capped and can be formed by ethoxylation of the respective alcohol with ethylene oxide. Also, methyl capped PEGs are commercially available from Union Catride under the trade name Methoxy Carbowax and from Aldrich Chemical Company under the name poly(ethylene glycol) methyl ether. These commercial methyl capped PEGs have molecular weights of 350 (n = about 7.5), 550 (n = about 12), 750 (n = about 16), 2000 (n = about 113). The capped PEG, or more typically lis alkali metal (Nat or K*) alkoxide, are reacted with a glycidyl ether to form the capping group used in preparing the polyesters of the present invention. See Flores-Gallardo et al, "Epoxy Ethers and Ether Amino Alcohols", J. Org. Chem., Vol. 12, (1947), pp 331-33, which describes a method for preparing dyictly! ether useful in the present invention.

A representative synthesis of one such capping group is as follows:

Step 1: 1-methoxy-2-hydroxy-3-chloropropane

Into a 2-I., three-necked, round bottom flask, fitted with a condenser, addition funnel, and magnetic stirrer were placed 730 ml (18 moles) of methanoal and 16.0 ml (0.25 moles) of methanesulfonic acid. To this refluxing mixture was added dropwise 496 ml (6.0 moles) of epichlorohydrin. The reaction mixture was stirred and refluxed for 18 hrs. After cooling to room temperature, 37.3 g (0.27 moles) of K₂CO₃ was added to the reaction mixture which was then stirred for 2 hrs. The filtrate was distilled at atmospheric pressure to remove methanol, then at reduced pressure (50 *-55 °C) to obtain 288 g (36% yield) of pro- duct.

The NMR spectrum of the product included the expected absence of epoxide resonances and the 40 addition of a methoxy resonance. The remaining two methylene, methine, and alcohol resonances were as expected for 1-methoxy-2-hydroxy-3-chloropropane.

Step 2: 1,2-epoxy-3-methoxypropane

Into a 2-1, three-necked, round bottom flask, fitted with a condenser, and mechanical stirrer were placed 200.2 g (1.6 moles) of the 1-methoxy-2-hydroxy-3-chloropropane from Step 1 and 1.6.1 of diethyl ether. The flask was immersed in an ice-water bath, and 96.0 g (2.4 moles) of NaOH was then added in small portions over a 2.5 hr. period to the vigorously stirred reaction mixture. The reaction mixture was allowed to warm to room temperature and was stirred overnight. The ether phase was then washed with H₂O (2 X 100 ml). The combined aqueous extracts were washed once with 200 ml of diethyl ether. The combined diethyl ether extracts were dired with Na₂SO_A. The dried extracts were distilled at atmospheric pressure to remove diethyl ether, then under reduced oressure (35° C) to obtain 39.5 a (67% vield) of produce.

The NMR spectrum of the product included the expected epoxide, methoxy and the methylene resonances for 1,2-epoxy-3-methoxypropane.

Step 3: Reaction of 1,2-epoxy-3 methoxypropane and poly(ethylene glycol) methyl ether to form capping group

Into a 250 ml, three-necked, round bottom flask, fitted with a condenser, addition funnel, and magnetic stirrer were placed 175.0 g (0.5 moles) of poly(ethylene glycol) methyl ether (M.W. 350) and 1.1 g (0.05 moles) of NaH. The mixture was stirred vigorously and heated to 80 °C under an argon atmosphere for 15 minutes. Then 88.4 g of the 1,2-epoxy-3-methoxy propane from Step 2 was added dropwise over a 6 ln. period. This reaction mixture was heated at 90 °C for 30 hrs. During this time, an additional 4.0 g can moles) of NaH was added periodically in small portions to maintain a pH of 10-11. (It is believed that some of the alkoxide generated by the NaH was consumed by a small amount of residual chlorosubstituted materials in the 1.2-epox-3-methoxypropane).

The reaction mixture was monitored by H-NMR and was considered complete following the absence of epoxide resonances. After 30 hours, the reaction mixture was allowed to cool to room temperature and 10.2 g (0.2 moles) of acetic acid was then added to neutralize the mixture. The reaction mixture was streed for 15 15 minutes, then excess acetic acid was removed on a Kugelrohr evaporator at 100°C for 5.5 hrs. This resulted in 255.1 g (978) yield of product.

The NMR spectrum of the product containing the capping group included the expected resonances for the methoxy and ethoxylate groups, and the absence of epoxide resonances.

Preferably, the only dicarboxylic acid used is terephthalic acid or its diester. However, minor amounts of 20 other aromatic dicarboxylic acids (or their diesters), or aliphatic dicarboxylic acids (or their diesters) can be included to the extent that the soil release properties are substantially maintained. Illustrative examples of other aromatic dicarboxylic acids which can be used include isophthalic acid, phthalic acid, naphthalene dicarboxylic acids, anthracene dicarboxylic acids, biphenyl dicarboxylic acids, oxydibenzoic acids, as well as mixtures of these acids. If aliphatic dicarboxylic acids are included, adpic, glutaric, succinic, 25 trimethyladipic, pimelic, azelaic, sebacic, suberic, 1.4-cyclohexane dicarboxylic acid and/or dodecanedicic acids can be used.

Illustrative examples of sulfonated aromatic dicarboxylic acids which can be used to prepare polyesters of the present invention include the alkyl metal salts of benzene-2,5-dicarboxy sulfonate, 2-naphthyl-dicarboxy-benzene sulfonate, phenyl-dicarboxy-benzene sulfonate, phenyl-dicarboxy-benzene sulfonate, phenyl-dicarboxy-benzene sulfonate, phenyl-dicarboxy-benzene sulfonate, 2,2-dimethyl phenyl-3, 5-dicarboxy benzenes sulfonate and phenyl-3, 5-dicarboxy-benzene sulfonate, preferred sulfonated salt is the 5-sulfoisophthalic acid sodium salt or its diester. Il branched polyesters are desired, a minor amount of a polycarboxylic acid (or its diester) selected from trimesic acid, trimellitic acid, and mixtures thereof can be used.

The preferred method for preparing block polyesters of the present invention comprises reacting the deeired mixture of lower dialkyl esters (methyl, ethyl, propyl or butyl) of the dicarboxylic acid with a mixture of the glycol (ethylene glycol), 1,2-propylene glycol or a mixture thereof) and the capping group. The glycol esters and oligomers produced in this seter interchange reaction are then polymerized to the desired degree. The ester interchange reaction are be conducted in accordance with reaction conditions generally used for ester interchange reactions. This ester interchange reaction is usually conducted at temperatures of from 120° to 220°C in the presence of an esterification catalyst. Alcohol is formed and constantly removed thus forcing the reaction to completion. The temperature and pressure of the reaction are desirably controlled so that glycol does not distill from the reaction mixture. Higher temperatures can be used if the reaction is conducted under pressure.

The catalysts used for the ester interchange reaction are those well known to the art. These catalysts include alkyl and alkaline earth metals, for example tithium, sodium, calcium, and magnesium, as well as transition and Group II B metals, for example antimony, maganese, cobalt, and zinc, usually as the respective oxides, carbonates, or acetates. Typically, antimony trioxide and calcium acetate are used.

The extent of the ester interchange reaction can be monitored by the amount of alcohol liberated or the disappearance of the disalkyl esters of the disasic acids in the reaction mixture as determined by high performance liquid chromatography (HPLC) or any other suitable method. The ester interchange reaction is desirably taken to more than 90% completion. Greater than 95% completion is preferred in order to decrease the amount of sublimates obtained in the polymerization step.

If desired, stabilizers such as phosphorus and phosphoric acid and esters thereof can be added at the end of the ester interchange step. The purpose of the stabilizier is to inhibit degradation, oxidation, and other side reactions; to destroy the catalytic activity of the ester interchange catalyst; and to prevent precipitation of insoluble metal carboxylates. Typically, stabilizers are not used to make the polyesters of the present invention.

When the ester interchange reaction is complete, the glycol ester products are then polymenzed to produce polywesters. The desired degree of polymerization can be determined by HPLC and ¹²C-NMR analysis. For commercial processes, the polymerization reaction is usually conducted at temperatures of from about 200° to about 280° C in the presence of a catalyst. Higher temperatures can be used but tend to produce darker colored products. Illustrative examples of catalysts useful for the polymerization step include antimony trioxide, germanium dioxide, titanium alkoxide, hydrated antimony pentoxide, and ester interchange catalysts such the as salts of zinc, cobalt, and maganese.

Excess glycol and other volatiles liberated during the reaction are removed under vacuum. The reaction is continued until polymerization is nearly complete based on analysis by ¹³C-NMR and/or reverse phase in HPLC and/or gel phase permeation. In addition to the desired polyesters, the crude composition obtained after synthesis contains starting reactants, as well as intermediate products.

Representative examples of specific block polyesters formed according to the present invention are as follows:

15 EXAMPLE 1

In a 250 mt, three-necked, round bottom flask, fitted with a magnetic stirrer and a modified claisen head (to support a condenser and receiver flask) is placed 10.6 g (0.17 moles) of ethylene glycol, and 0.4 g (0.5% w/v) of Sb₂O₂ catalyst. This mixture is heated to 150°C for 1 hr. to predissolve the catalyst and then 20 cooled to room temperature. Then, 49.9 g (0.095 moles of the capping group (reaction product of 1.2 epoxy-3-methoxy propane and poly(ethyleneglycol) methyl ether of M.W. 350, 252 g (0.13 moles) of dimethyl terephthalate, and 0.1 g (0.1% w/v) of butylated hydroxyloluene is added. The reaction mixture is heated under argon to 180°C over a 2 hr. period and then held at 180°C for 23 hrs. During this time, 8.0 g (95% of theoretical value) of methanol is distilled from the reaction mixture. The reaction mixture is then 20 collect to room temperature, placed in a vacuum, heated to 200°C over a period of 1 hr., and then held at 200°C for 5 hrs. The reaction is determined to be complete by using ¹⁰C-MMR.

EXAMPLE 2

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A polyester is prepared by substituting 1,2-propylene glycol for ethylene glycol, using reaction conditions similar to those of Example 1.

Method for Determining Degree of Polymerization

3s A method for determining the degree of polymerization of the polyesters of the present invention involves: (1) alcohol fractionation of the crude polyester composition obtained after synthesis: (2) high performance liquid chromatographic (HPLC) separation of the methanol soluble fraction to yield additional fractions; and (3) ¹²C NMR analysis to determine backbone length (i.e. value for u) of the polyesters present in each of the various HPLC fractions. See U.S. application Serial No. 801,020 to Eugene P. Gosselink, filed November 22, 1995, pages 18-23, which provides a detailed description of this method for determining the degree of polymerization of similar polyesters.

DETERGENT COMPOSITIONS

45 Soil Release Component

The compounds of the present invention are particularly useful in detergent compositions to provide soil release properties. These compositions can be used as laundry detergents, laundry additives, and laundry pre-treatments.

The detergent compositions of the present invention comprise a soil release component which contains an effective amount of the soil release compounds previously defined. What is an "effective amount" will depend upon the particular soil release compounds used, the particular type of detergent formulation (liquid, granular, etc.) and the benefits desired. Usually, the soil release compounds are effective when included fin an amount from about 0.01 to about 10% by weight of the composition. In terms of soil release benefits, specifered detergent compositions can comprise from about 0.1 to about 5% by weight of the soil release compounds, but typically comprise from about 0.3 to about 3% by weight of these compounds.

For granular detergent formulations, the soil release component typically comprises the soil release compounds, plus any protective enrobing material. In making granular detergent formulations, the soil

release compounds could be exposed to highly alkaline materials such as NaOH and KOH. The soil release compounds, in particular those having shorter backbones, can be degraded by alkaline environments, especially those above a pH of about 8.5. Accordingly, the soil release compounds are preferably enrobed in a material which protects them from the alkaline environment of a granular detergent formulation yet permits the soil release compounds to be dispersed in the laundering operation.

Suitable enrobing materials include the nonionic surfactants, polyethylene glycols (PEG), fatty acids, fatty acid selser of alcohols, diols and polyols, anionic surfactants, film forming polymers and mixtures of these materials. Examples of suitable nonionic surfactant enrobing materials are described in the Deteregent Surfactant section of this application. Examples of suitable PEG enrobing materials are those having an average M.W. of from about 2,000 to 15,000, preferably from about 3,000 to about 10,000 and most preferably from about 4,000 to about 8,000. Examples of suitable tatty acid enrobing materials are the higher fatty acids having from 12 to 18 carbon atoms. Examples of suitable fatty acid seter enrobing materials include the sorbitant ratty acid esters (e.g. sorbitan mondaurate). Other examples of suitable enrobing materials, including anionic surfactants and film forming polymers, are disclosed in U.S. Patent 14, 4486,327 to Murphy et al., issued December 4, 1984. The soil release compounds can be enrobed according to the methods disclosed in this Murphy et al. patent.

For līquid detergent formulations, the soil release component can be comprised entirely of soil release compounds or can further include a water-soluble organic solvent or an hydrotrope to aid in dissolving the soil release compounds. Suitable organic solvents are usually aromatic and can include ethyl benzoate, properties of the process of the proc

28 The amount, or oven need for, organic solvents or hydrotropes to prepare liquid detergent formulations containing the soil release compounds of the present invention will depend upon the compounds used, especially what fraction thereof is water soluble, the ingredients present in the laundry detergent system, and whether an isotropic, hornogeneous liquid is desired. For isotropic liquid detergent formulations, the soil release compounds need to be dissolved as much as possible which sometimes requires the use of organic solvents or hydrotropes. Also, it has been found that dissolving the compounds in the liquid detergent formulations makes them more effective as soil release capents.

Besides organic solvents or hydrotropes, greater amounts of water-soluble soil release compounds ons.

For example, soil release component to ad in the preparation of isotropic liquid detergent formulations.

For example, soil release polyesters with backbones having from about 3 to about 5 (u = about 3 to about 3 5 12-proylene terphthalate units and having a capping group at each end based on a methyl capped PEG (M.W. 350) and a methoxy glycidyl ether reaction product are water-soluble. In addition, soil release polyesters prepared from dimethyl terephthalate, 1/2-propylene glycol and the capping group typically contain a substantial fraction of water-soluble polyesters (both active and inactive types) which aid in dissolving less water-soluble soil release polyesters in the liquid detergent formulation. Partial or total inclusion of 1.2-propylene glycol generally increases the solubility of the soil release polyesters. The more water-soluble 1.2-propylene glycol dased soil release polyesters are particularly useful in making isotropic liquid detergent formulations which have a large number of ingredients and low water content.

Detergent Surfactants

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The amount of detergent surfactant included in the detergent compositions of the present invention can vary from about 1 to about 75% by weight of the composition depending upon the detergent surfactant(s) used, the type of composition to be formulated (e.g. granular, liquid) and the effects desired. Preferably, the detergent surfactant(s) comprises from about 10 to about 50% by weight of the composition, and most preferably from about 15 to about 40% by weight. The detergent surfactant can be nonlonic, anionic, ampholytic, writeriorinc, catomic, or a mixture thereof:

A. Nonionic Surfactants

Suitable nonionic surfactants for use in detergent compositions of the present invention are generally disclosed in U.S. Patent 3,99,678 to Laughlin et al., issued December 30, 1975 at column 13, line 14 through column 16, line 6. Classes of nonionic surfactants included are:

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1. The polyethylene oxide condensates of alkyl phenols. These compounds include the condensation products of alkyl phenols having an alkyl group containing from about 6 to 12 carbon atoms in either a straight chain or branched chain configuration with ethylene oxide, the ethylene oxide being present in an amount equal to 5 to 25 moles of ethylene oxide per mole of alkyl phenol. The alkyl substituent in such compounds can be derived, for example, from polymerized propylene, diisobutylene. Examples of compounds of this type include nonyl phenol condensed with about 15 moles of ethylene oxide per mole of nonyl phenol; dodecylphenol condensed with about 12 moles of ethylene oxide per mole of phenol; dinonyl phenol condensed with about 15 moles of ethylene oxide per mole of phenol; and diisooctyl phenol condensed with about 15 moles of ethylene oxide per mole of phenol; and diisooctyl phenol condensed with about 15 moles of ethylene oxide per mole of phenol. Commercially available nonlonic surfactants of this type include Igepal CD-630, marketed by the GAF Corporation, and Triton X-45, X-114, X-100, and X-102, all marketed by the GAF Company.

2. The condensation products of alighatic alcohols with from about 1 to about 25 moles of ethylene oxide. The aliqy chain of the alightatic alcohol can either be straight or branched, primary or secondary, and generally contains from about 8 to about 22 carbon atoms. Examples of such ethoxylated alcohols include the condensation product of about 9 moles of ethylene oxide per mole of alcohols, and the condensation product of about 9 moles of ethylene oxide with coconut alcohol (a mixture of fatty alcohols with alkyl chains varying in length from 10 to 14 carbon atoms). Examples of commercially available nonionic surfactants of this type include Terigitol 15-89, marketed by Union Carbide Corporation, Neodol 45-9, Neodol 23-6.5, Neodol 45-7, and Neodol 45-4, marketed by Shell Chemical Company, and Kyro EOB. marketed by The Procter & Gambie Company.

3. The condensation products of ethylene oxide with a hydrophobic base formed by the condensation of propylene oxide with propylene glycol. The hydrophobic portion of these compounds has a molecular weight of from about 1500 to 1800 and exhibits water insolubility. The addition of polyoxyethylene moieties to this hydrophobic portion tends to increase the water solubility of the molecule as a whole, and the liquid character of the product is retained up to the point where the polyoxyethylene content is about 50% of the total weight of the condensation product, which corresponds to condensation with up to about 40 moles of ethylene oxide. Examples of compounds of this type include certain of the commerciality available Pluronic surfactants, marketed by Wyandotte Chemical Corporation.

4. The condensation products of ethylene oxide with the product resulting from the reaction of propylene oxide and ethylenediamine. The hydrophobic molety of these products consists of the reaction product of ethylenediamine and excess propylene oxide, the molety having a molecular weight of from about 2500 to about 3000. This hydrophobic molety is condensed with ethylene oxide to the extent that the condensation product contains from about 40% to about 80% by weight of polyoxyethylene and has a molecular weight of from about 5,000 to about 11,000. Examples of this type of nonionic surfactant include certain of the commercially available Tetronic compounds, marketed by Wyandotte Chemical Corporation.

5. Semi-polar nonionic detergent surfactants which include water-soluble amine oxides containing one alkyl moiety of from about 10 to 18 carbon atoms and 2 moleties selected from the group consisting of alkyl groups and hydroxyalkyl groups containing from 1 to about 3 carbon atoms; water-soluble phosphine oxides containing one alkyl molety of from about 10 to 18 carbon atoms and 2 moleties selected from the group consisting of alkyl groups and hydroxyalkyl groups containing from about 1 to 3 carbon atoms; and water-soluble sulfoxides containing one alkyl molety of from about 10 to 18 carbon atoms and a moiety selected from the group consisting of alkyl and hydroxyalkyl moieties of from about 1 to 3 carbon atoms.

Preferred semi-polar nonionic detergent surfactants are the amine oxide detergent surfactants having the formula

wherein \mathbb{R}^3 is an allyl, hydroxyalkyl, or alkyl phenyl group or mixtures thereof containing from about 8 to about 25 carbon atoms; \mathbb{R}^4 is an alkylene or hydroxyalkylene group containing from 2 to 3 carbon atoms or mixtures thereof; x is from 0 to about 3; and each \mathbb{R}^3 is an alkyl or hydroxyalkyl group containing from 1 to about 3 carbon atoms or a polyethylene oxide group containing from one to about 3 ethylene oxide groups. The \mathbb{R}^6 groups can be attached to each other, e.g., through an oxygen or nitrogen atom to form

a ring structure.

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Preferred amine oxide detergent surfactants are C₁₀-C₁₈ alkyl dimethyl amine oxide and C₈-C₁₂ alkoxy ethyl dihydroxy ethyl amine oxide.

6. Alkylpolysaccharides disclosed in European Patent Application 70,074 to Ramon A. Llenado, published January 19, 1983, having a hydrophobic group containing from about 6 to about 30 carbon atoms, preferably from about 10 about 16 carbon atoms and a polysaccharide, e.g., a polyplycoside, hydrophilic group containing from about 1½ to about 10, preferably from about 1½ to about 3, most preferably from about 16 about 27 saccharide units. Any reducing saccharide containing 5 or 6 carbon atoms can be used, e.g. glucose, galactose and galactosyl moieties can be substituted for the glucosyl moieties (Optionally the hydrophobic group is attached at the 2, 3, 4, etc. positions thus giving a glucose or galactose as opposed to a glucoside or galactoside). The intersaccharide bonds can be, e.g., between the one position of the additional saccharide units and the 2-, 3-, 4-, and/or 6 positions on the preceding saccharide units.

Öptionally, and less desirably, there can be a polyalkyleneoxide chain joining the hydrophobic molety and the polysaccharide molety. The preferred alkyleneoxide is ethylene oxide. Typical hydrophobic groups include alkyl groups, either saturated or unsaturated, branched or unbranched containing from about 8 to about 16, preferably from about 10 to about 16, carbon atoms. Preferably, the alkyl group is a straight chain saturated alkyl group. The alkyl group can contain up to 3 hydroxy groups and/or the polyalkyleneoxide chain can contain up to about 10, preferably less than 5, most preferably 0, alkyleneoxide moleties. Suitable alkyl polysaccharides are octyl, nonyldecyl, nondecyllodecyl, tridecyl, tetradecyl, pentadecyl, heptadecyl, and octadecyl, di-, tri-, tetra-, penta-, and hexaglucosides, galactosides, factosides, glucoses, fructosides, fructoses, and/or galactoses. Suitable mixtures include coconut alkyl, di-, tri-, tetra-, and pentadulcosides and tallow alkyl tetra-, penta-, and hexaglucosides.

The preferred alkylpolyglycosides have the formula

R2O(C_nH_{2n}O)_t(alvcosvI)_v

wherein R² is selected from the group consisting of alkyl, alkylphenyl, hydroxyalkylphenyl, and mixtures thereof in which the alkyl groups contain from about 10 to about 10, preferably from about 12 to about 14, carbon atoms; n is 2 or 3, preferably 2; t is from 0 to about 10, preferably 0; and x is from 1½ to about 10, preferably 0; and x is from 1½ to about 10, preferably from about 1½ to about 3, most preferably from about 1.6 to about 2.7. The glycosyl is preferably derived from glucoss. To prepare these compounds, the alcohol or alkyl polyethoxy alcohol is formed first and then reacted with glucose, or a source of glucose, to form the glucoside (attachment at the 1-position). The additional glycosyl units can then be attached between their 1-position and the preceding glycosyl units 2-, 3-, 4- and/or 6- position, preferably predominately the 2-position.

7. Fatty acid amide detergent surfactants having the formula:

wherein R^6 is an alkyl group containing from about 7 to about 21 (preferably from about 9 to about 17) carbon atoms and each R^7 is selected from the group consisting of hydrogen, C₁-C₄ alkyl, C₁-C₅ hydroxyalkyl, and -(C₂H₄O)₃H where x varies from about 1 to about 3. Preferred amides are C₃-C₂₀ ammonia amides, monoethanolamides, diethanolamides, and isopropanol amides.

B. Anionic Surfactants

Anionic surfactants suitable in detergent compositions of the present invention are generally disclosed in U.S. Patent 3,929,978 to Laughlin et al., issued December 30, 1975 at column 23, line 58 through column 29, line 23. Classes of anionic surfactants included are:

- Ordinary alkali metal soaps such as the sodium, potassium, ammonium and alkylolammonium salts of higher fatty acids containing from about 8 to about 24 carbon atoms, preferably from about 10 to about 20 carbon atoms.
 - 2. Water-soluble salts, preferably the alkali metal, ammonium and alkylolammonium salts, of organic sulfuric reaction products having in their molecular structure an alkyl group containing from about 10 to

about 20 carbon atoms and a sulfonic acid or sulfuric acid ester group. (Included in the term "alkyl" is the alkyl portion of acyl groups.)

Examples of this group of anionic surfactants are the sodium and potassium alkyl sulfates, especially those obtained by sulfating the higher alcohols (Ga-Cu₀ carbon atoms) such as those produced by reducing the glycerides of tallow or occonut oil; and the sodium and potassium alkylbenzene sulfonates in which the alkyl group contains from about 9 to about 15 carbon atoms, in straight chain or branched chain configuration, e.g., those of the type described in U.S. Patents 2200.099 and 2.477.383. Especially valuable are linear straight chain alkylbenzene sulfonates in which the average number of carbon atoms in the alkyl group is from about 11 to 13, abbreviated as C₁1-C₁1AS.

Preferred anionic surfactants of this type are the alkyl polyethoxylate sulfates, particularly those in which the alkyl group contains from about 10 to about 22, preferably from about 12 to about 18 carbon atoms, and wherein the polyethoxylate chain contains from about 1 to about 15 ethoxylate moieties preferably from about 1 to about 3 ethoxylate moieties. These anionic detergent surfactants are particularly desirable for formulation heavy-duxt liquid leaundy detergent compositions.

Other anionic surfactants of this type include sodium alkyl glyceryl ether sulfonates, especially those ethers of higher alcohols derived from tallow and occonut oil; sodium coconut oil fatty acid monoglyceride sulfonates and sulfates; sodium or potassium salts of alkyl phenol ethylene oxide ether sulfates containing from about 1 to about 10 units of ethylene oxide per molecule and wherein the alkyl groups contain from about 8 to about 12 carbon atoms; and sodium or potassium salts of alkyl ethylene oxide ether sulfates containing about 1 to about 10 units of ethylene oxide per molecule and wherein the alkyl group contains from about 10 to about 20 carbon atoms.

Also included are water-soluble salts of esters of alpha-sulfonated fatty acids containing from about 8 to 20 carbon atoms in the fatty acid group and from about 1 to 10 carbon atoms in the ester group; water-soluble salts of 2-acytoxy-alkane-1-sulfonic acids containing from about 2 to 9 carbon atoms in the acyl group and from about 9 to about 23 carbon atoms in the alkane moiety; alkyl either sulfates containing from about 10 to 20 carbon atoms in the alkyl group and from about 1 to 30 moles of ethylene oxide; water-soluble salts of olefin sulfonates containing from about 12 to 24 carbon atoms; and beta-alkylovy alkane sulfonates containing from about 1 to 30 carbon atoms in the alkyl group and from about 8 to 20 carbon atoms in the alkylan moiety.

- Anionic phosphate surfactants.
 - 4. N-alkyl substituted succinamates.

C. Ampholytic Surfactants

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35 Ampholytic surfactants can be broadly described as aliphatic derivatives of secondary or tertiary amines, or aliphatic derivatives of heterocyclic secondary and tertiary amines in which the aliphatic radical can be straight chain or branched and wherein one of the aliphatic substituents contains from about 8 to 18 carbon atoms and at least one contains an anionic water-solubilizing group, e.g. carboxy, sulfonate, sulfate. See U.S Patienti 3,929,678 to Lauphlin et al., issued December 30, 1975 at column 19, lines 18-35 for examples of ampholytic surfactants.

D. Zwitterionic Surfactants

Zwitterionic surfactants can be broadly described as derivatives of secondary and teriary amines, derivatives of heterocyclic secondary and teriary amines, or derivatives of quaternary ammonium, quaternary phosphonium or teriary sulfonium compounds. See U.S. Patent 3,929,678 to Laughlin et al., issued December 30, 1975 at column 19, line 38 through column 22, line 48 for examples of zwitterionic surfactants.

50 E. Cationic Surfactants

Cationic surfactants can also be included in detergent compositions of the present invention. Suitable cationic surfactants include the quaternary ammonium surfactants having the formula:

55 [R²(OR³)_v][R⁴(OR³)_v]₂ R⁵ N+X[−]

wherein R² is an alkyl or alkyl benzyl group having from about 8 to about 18 carbon atoms in the alkyl chain; each R³ is selected from the group consisting of -CH₂CH₂-, -CH₂CH(CH₃)-, -CH₂CH(CH₂OH)-,

-CH₂CH₃CH₃-C, and mixtures thereof; each R⁴ is selected from the group consisting of C₁-C₄ alkyl, C₁-C₄ hydroxyalkyl, benzyl, ring structures formed by joining the two R¹ groups, CH₂CHOHCHOHCOHCH₂OH wherein R⁶ is any hexose or hexose polymer having a molecular weight less than about 1000, and hydrogen when y is not 0; R⁶ is the same as R⁴ or is an alkyl chain 5 wherein the total number of carbon atoms of R⁷ plus R⁶ is not more than about 10; each y is from 0 to about 10 and the sum of the y values is from 0 to about 10 and the sum of the y values is from 0 to about 10 and the sum of the y values is from 0 to about 10 and the sum of the y values is from 0 to about 10 and the sum of the yalues is from 0 to about 10 and the sum of the yalues is from 0 to about 10 and the sum of the yalues is from 0 to about 10 and the sum of the yalues is from 0 to about 10 and the sum of the yalues is from 0 to about 10 and the sum of the yalues is from 0 to about 10 and the sum of the yalues is from 0 to about 10 and the yalues is from 0 to about 10 and the yalues is from 0 to about 10 and the yalues is from 0 to about 10 and the yalues is from 0 to about 10 and the yalues is from 0 to about 10 and the yalues is from 0 to about 10 and 10 and

Preferred of the above are the alkyl quaternary ammonium surfactants, especially the mono-long chain alkyl surfactants described in the above formula when RP is selected from the same groups as R¹. The most preferred quaternary ammonium surfactants are the chloride, bromide and methylsulfate $C_0 - C_{1:0}$ alkyl to trimethylammonium salts, $C_0 - C_{1:0}$ alkyl di(flydroxyelthylmethylammonium salts, the $C_0 - C_{1:0}$ alkyl hydroxyelthylmethylammonium salts, and $C_0 - C_{1:0}$ alkyloxypropyl trimethylammonium salts, and $C_0 - C_{1:0}$ alkyloxypropyl trimethylammonium methylsulfate, lauryl trimethylammonium chloride, myristyl trimethylammonium bromide and coconut trimethylammonium chloride and methylsulfate are particularly preferred.

Other useful cationic surfactants are disclosed in U.S. Patent 4,259,217 to Murphy, issued March 31, 15, 1981.

Detergent Builders

Detergent compositions of the present invention can optionally comprise inorganic or organic detergent builders to assist in mineral hardness control. When included, these builders typically comprise up to about 60% by weight of the detergent composition. Built liquid formulations preferably comprise from about 1 to about 25% by weight detergent builder, most preferably from about 3 to about 20% by weight, while built granular formulations preferably comprise from about 5 to about 50% by weight detergent builder, most preferably from about 10 to about 30% by weight.

Suitable detergent builders include crystalline aluminosilicate ion exchange materials having the formula:

Na_z[(A1O₂)_z • (SiO₂)_v • xH₂O

30 wherein z and y are at least about 6, the mole ratio of z to y is from about 1.0 to about 0.5; and x is from about 10 to about 264. Amorphous hydrated aluminosilicate materials useful herein have the empirical formula

 $M_z(zA1O_2 \cdot ySiO_2)$

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wherein M is sodium, potassium, ammonium or substituted ammonium, z is from about 0.5 to about 2; and y is 1; this material having a magnesium ion exchange capacity of at least about 50 milligram equivalents of CaCO₂ hardness per gram of anhydrous aluminosilicate.

The aluminosilicate ion exchange builder materials are in hydrated form and contain from about 10% to 40 about 28% of water by weight if crystalline, and potentially even higher amounts of water if amorphous. Highly preferred crystalline aluminosilicate ion exchange materials contain from about 18% to about 22% water in their crystal matrix. The preferred crystalline aluminosilicate ion exchange materials are further characterized by a particle size diameter of from about 0.1 micron to about 10 microns. Amorphous materials are often smaller, e.g., down to less than about 0.01 micron. More preferred ion exchange 45 materials have a particle size diameter of from about 0.2 micron to about 4 microns. The term "particle size diameter" represents the average particle size diameter of a given ion exchange material as determined by conventional analytical techniques such as, for example, microscopic determination utilizing a scanning electron microscope. The crystalline aluminosilicate ion exchange materials are usually further characterized by their calcium ion exchange capacity, which is at least about 200 mg. equivalent of CaCO3 water 50 hardness/g, of aluminosilicate, calculated on an anhydrous basis, and which generally is in the range of from about 300 mg, eq./g, to about 352 mg, eq./g. The aluminosilicate ion exchange materials are still further characterized by their calcium ion exchange rate which is at least about 2 grains Ca++/gallon/minute/gram/gallon of aluminosilicate (anhydrous basis), and generally lies within the range of from about 2 grains/gallon/minute/gram/gallon to about 6 grains/gallon/minute/gram/gallon, based on calcium 55 ion hardness. Optimum aluminosilicates for builder purposes exhibit a calcium ion exchange rate of at least about 4 grains/gallon/minute/gram/gallon.

The amorphous aluminosilicate ion exchange materials usually have a Mg++ exchange capacity of at least about 50 mg, eq. CaCO₃/g, (12 mg, Mg++/g,) and a Mg++ exchange rate of at least about 1

grain/gallon/minute/gram/gallon. Amorphous materials do not exhibit an observable diffraction pattern when examined by Cu radiation (1.54 Angstrom Units).

Useful aluminosilicate ion exchange materials are commercially available. These aluminosilicates can be crystalline or amorphous in structure and can be naturally-occurring aluminosilicates or synthetically derived. A method for producing aluminosilicate ion exchange materials is disclosed in U.S. Patent 3,995,669 to Krummel, et al. issued October 12, 1976. Preferred synthetic crystalline aluminosilicate ion exchange materials useful heroria ner available under the designations 20otlot A, Zeolitic P (B), and Zeolitic A. In an especially preferred embodiment, the crystalline aluminosilicate ion exchange material has the formula

Na₁₂[(A1O₂)₁₂(SiO₂)₁₂]•xH₂O

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wherein x is from about 20 to about 30, especially about 27.

Other examples of detergency builders include the various water-soluble, alkali metal, ammonium or 5 substituted ammonium phosphates, polyphosphates, phosphonates, polyphosphonates, carbonates, silicates, borates, polyhydroxysulfonates, polyacetates, carboxylates, and polycarboxylates. Preferred are the alkali metal, especially sodium, salts of the above.

Specific examples of inorganic phosphate builders are sodium and potassium tripolyphosphate, pyrophosphate, polyphosphate, polyphosphate, prophosphate, prophosphate, prophosphate, prophosphate, Examples of polyphosphonate builders are the sodium and potassium salts of ethylene-1,1-diphosphonic acid, the sodium and potassium salts of ethane 1-hydroxy-1,1-diphosphonic acid and the sodium and potassium salts of ethane, 1,1,2-triphosphonic acid. Other phosphorus builder compounds are disclosed in U.S. Patents 3,159,561; 3,213,030; 3,422,017; 3,420,176 and 3,400,148.

Examples of nonphosphorus, inorganic builders are sodium and potassium carbonate, bicarbonate, sesquicarbonate, tetraborate decahydrate, and silicate having a mole ratio of SIO₂ to alkali metal oxide of from about 10.5 to about 4.4 do. referably from about 10.0 to about 2.4.

Useful water-soluble, nonphosphorus organic builders include the various alkali metal, ammonium and substituted ammonium polyacetates, carboxylates, polycarboxylates and polyhydroxysulfonates. Examples of polyacetate and polycarboxylate builders are the sodium, potassium, lithium, ammonium and substituted 30 ammonium salts of ethylenediamine tetraacetic acid, nitrilotriacetic acid, oxydisuccinic acid, mellitic acid, benzene polycarboxylic acids, citric acid, and 2-hydroxyethyl othylenediamine triacetic acid.

Highly preferred polycarboxylate builders are disclosed in U.S. Patent No. 3,308,067 to Diehl, issued March 7, 1967. Such materials include the water-soluble salts of homo- and copolymers of aliphatic carboxylic acids such as maleic acid, itaconic acid, mesaconic acid, fumaric acid, aconitic acid, citraconic acid and methylenemalonic acid.

Other builders include the carboxylated carbohydrates disclosed in U.S. Patent 3,723,322 to Diehl issued March 28, 1973.

Other useful builders are sodium and potassium carboxymethyloxymalonate, carboxymethyloxysuccinate, cis-cyclohexanehexacarboxylate, cis-cyclopentaneletracarboxylate phloroglucinol trisulfonate, watersoluble polyacrylates (having molecular weights of from about 2,000 to about 200,000 for example), and the copolymers of maleic anhydride with vinyl methyl ether or ethylene.

Other suitable polycarboxylates are the polyacotal carboxylates disclosed in U.S. Patent 4,144,226, to Crutchfield et al. issued March 13, 1979, and U.S. Patent 4,244,495, to Crutchfield et al., issued March 27, 1979. These polyacetal carboxylates can be prepared by bringing together under polymerization conditions an ester of glyoxylic acid and a polymerization initiator. The resulting polyacetal carboxylate ester is then attached to chemically stable end groups to stabilize the polyacetal carboxylate against rapid depolymerization in alkaline solution, converted to the corresponding salt, and added to a surfactant.

Clay Soil Removal/Anti-Redeposition Agents

Laundry detergent compositions of the present invention desirably include a clay soil removal and/or anti-redeposition agent. These clay soil removal/anti-redeposition agents are usually included at from about 0.1 to about 10% by weight of the composition. In terms of the benefits achieved, preferred detergent compositions can comprise from about 0.5 to about 5% by weight of these agents. Typically, these preferred compositions comprise from about 1 to about 3% by weight of these agents.

One group of preferred clay soil removal/anti-redeposition agents are the ethoxylated amines disclosed in European patent application 112,593 to James M. Vander Meer, published July 4, 1984. These ethoxylated amines are selected from the group consisting of:

(1) ethoxylated monoamines having the formula:

(2) ethoxylated diamines having the formula:

$$(R^2)_2 - N - R^1 - N - (R^2)_2$$

or

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(3) ethoxylated polyamines having the formula:

$$R^3 - [(A^1)_{q} - (R^4)_{t} - N - L - X]_{p}$$

(4) ethoxylated amine polymers having the general formula:

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(5) mixtures thereof; wherein A1 is

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or -O-:

R is H or C₁-C₁ alkyl or hydroxyalkyl; R¹ is C₂-C₁₂ alkylene, hydroxyalkylene, alkenylene, arylene or alkenylene, or a C₂-C₂ oxyalkylene moiety having from 2 to about 20 oxyalkylene units provided that no O-N bonds are formed; each R² is C₁-C₂ alkyl or hydroxyalkyl, the moiety -L-X, or two R² bogother form the moiety -(CH₂)_xA²-(CH₂)_x, wherein A² is -O- or -CH₂-, r is 1 or 2, s is 1 or 2, and r + s is 3 or 4; X is a nonionic group, an anionic group or mixture thereof; R³ is a substituted C₂-C₁₂ alkyl, hydroxyalkyl, alkenyl, aryl, or alkaryl group having p substitution sites; R¹ is C₁-C₁₂ alkylene, hydroxyalkylene.

alkenylene, arylene or alkarylene, or a C2-C3 oxyalkylene moiety having from 2 to about 20 oxyalkylene units provided that no O-O or O-N bonds are formed; L is a hydrophilic chain which contains the polyoxyalkylene mojety f(R5O), (CH2CH2O), h, wherein R5 is C2-C4 alkylene or hydroxyalkylene and m and n are numbers such that the moiety -(CH2CH2O)- comprises at least about 50% by weight of said polyoxyalkylene moiety; for said monoamines, m is from 0 to about 4, and n is at least about 12; for said diamines, m is from 0 to about 3, and n is at least about 6 when R1 is C2-C3 alkylene, hydroxyalkylene. or alkenylene, and at least about 3 when R1 is other than C2-C3 alkylene, hydroxyalkylene or alkenylene; for said polyamines and amine polymers, m is from 0 to about 10 and n is at least about 3; p is from 3 to 8; q is 1 or 0; t is 1 or 0, provided that t is 1 when q is 1; w is 1 or 0; x + y + z is at least 2; and y +

Another group of preferred clay soil removal/anti-redeposition agents are the cationic compounds disclosed in European patent application 111.965 to Young S. Oh and Eugene P. Gosselink, published June 27, 1984. These cationic compounds are selected from the group consisting of:

(1) ethoxylated cationic monoamines having the formula:

$$R^2 - \frac{R^2}{N^4} - L - X$$

(2) ethoxylated cationic diamines having the formula:

or

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$$(x-L-)_2-M^2-R^1-M^2-R^2$$

wherein M1 is an N+ or N group; each M2 is an N+ or N group, and at least one M2 is an N+ group; (3) ethoxylated cationic polyamines having the formula:

$$R^4 - [(A^1)_q - (R^5)_t - M^2_{R^2} - L - X]_p$$

- (4) ethoxylated cationic polymers which comprise a polymer backbone, at least 2 M groups and at least 50 one L-X group, wherein M is a cationic group attached to or integral with the backbone and contains an N+ positively charged center; and L connects groups M and X or connects group X to the polymer backbone: and
 - (5) mixtures thereof;
- 55 wherein A1 is

or -O-, R is H or C1-C4 alkyl or hydroxyalkyl, R1 is C2-C12 alkylene, hydroxyalkylene, alkenylene, arylene or alkarylene, or a C2-C3 oxyalkylene moiety having from 2 to about 20 oxyalkylene units provided that no O-N bonds are formed; each R2 is C1-C4 alkyl or hydroxyalkyl, the moiety -L-X or two R2 together form the moiety -(CH₂),-A²-(CH₂),-, wherein A² is -O- or -CH₂-, r is 1 or 2, s is 1 or 2 and r + s is 3 or 4; each R3 is C1-C8 alkyl or hydroxyalkyl, benzyl, the moiety -L-X, or two R3 or one R2 and one R3 together form the moiety -(CH2)r-A2-(CH2)s-; R4 is a substituted C3-C12 alkyl, hydroxyalkyl, alkenyl, aryl or alkaryl group having p substitution sites; R5 is C1-C12 alkylene, hydroxyalkylene, alkenylene, arylene or alkarylene, or a C2-C3 oxyalkylenemoiety having from 2 to about 20 oxyalkylene units provided that no O-O or O-N bonds are formed; X is a nonionic group selected from the group consisting of H, C1-C4 alkyl or hydroxyalkyl ester or ether groups, and mixtures thereof; L is a hydrophilic chain which contains the polyoxyalkylene moiety f(R6O)m(CH2CH2O)n1; wherein R6 is C3-C4 alkylene or hydroxyalkylene and m and n are numbers such that the moiety -(CH2CH2O)- comprises at least about 50% by weight of said polyoxyalkylene moiety; d is 1 when M2 is N+ and is 0 when M2 is N; n is at least about 12 for said cationic monoamines, is at least about 6 for said cationic diamines and is at least about 3 for said cationic polyamines and cationic polymers; p is from 3 to 8; q is 1 or 0; and t is 1 or 0, provided that t is 1 when a is 1.

Other clay soil removal/anti-redeposition agents which can be used include the ethoxylated amine polymers disclosed in European patent application 111,984 to Eugene P. Gosselink, published June 27, 1984; the zwitterionic compounds disclosed in European patent application 111,976 to Donn N. Rubingh and Eugene P. Gosselink, published June 27, 1984; the zwitterionic polymers disclosed in European patent application 112,592 to Eugene P. Gosselink, published July 4, 1984; and the amine oxides disclosed in U.S. 35 Patent 4,548,744 to Connor, issued October 22, 1985.

Other Optional Detergent Ingredients

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Other optional ingredients which can be included in detergent compositions of the present invention, in 40 their conventional art-established levels for use (i.e., from 0 to about 20%), include solvents, bleaching agents, bleach activators, other soil-suspending agents, corrosion inhibitors, dyes, fillers, optical brighteners, germicides, pH adjusting agents (monoethanolamine, sodium carbonate, sodium hydroxide), enzymes, enzyme-stabilizing agents, perfumes, fabric softening components, static control agents,

45 General Detergent Formulations

Except for the previously described enrobing of the soil release compound, granular formulations embodying the detergent compositions of the present invention can be formed by conventional techniques, i.e., by slurrying the individual components in water and then atomizing and spray-drying the resultant mixture, or by pan or drum granulation of the ingredients. Granular formulations preferably comprise from about 10 to about 30% detergent surfactant, usually anionic, and most preferably about 15 to about 25%

Liquid formulations embodying the detergent compositions can be built or unbuilt. If unbuilt, these compositions conventionally contain approximately 15 to 50% (preferably 20 to 35%) total surfactant, from 0 to 5% (preferably from 0 to 2%) of an organic base such as a mono-, di-, or tri-alkanol amine, a neutralization system such as an alkali metal hydroxide and a lower primary alcohol such as ethanol or isopropanol, and approximately 20 to 80% water.

Built liquid detergent compositions can be in the form of single phase liquids provided that the builder is solubilized in the mixture at its level of use. Such liquids conventionally contain 10 to 40% (preferably 15 to 25%) builder which can be organic or inorganic, up to 10% of a hydrotrope system, and 20 to 80% water. Built liquid detergents incorporating components that of orm heterogeneous mixtures of relevate of builder that cannot be completely dissolved) can also comprise detergent compositions of the present invention. Such liquids conventionally employ viscosity modifiers to produce systems having plastic shear characteristics to maintain stable dispersions and to prevent phase separation or solid settlement. Care should also be taken to avoid exposing the soil release compounds to highly alkaline environments, e.g. those above a pH of about 8.5, during processing of the liquid detergent 10 formulation.

A description of some prefered detergent formulations is as follows:

A. Near Neutral Wash pH Detergent Formulations

While the detergent compositions of the present invention are operative within a wide range of wash pHs, they are particularly suitable when formulated to provide a near neutral wash pH. i.e. an initial pH of from about 6.0 to about 8.5 at a concentration of from about 0.1 to about 2% by weight in water at 20 °C. Near neutral wash pH formulations are better for enzyme statishilty and for preventing stains from settings. The near neutral pH of such formulations is also desirable to insure long term activity for the soil release compounds, especially those having shorter backbones. In such formulations, the product pH is preferably from about 7.5 to about 8.0.

Preferred near neutral wash pH detergent formulations are disclosed in U.S. Patent 4,561,998 to Wertz et al. issued December 31, 1985. These preferred formulations comprise:

 (a) from about 2 to about 60% (preferably from about 10 to about 25%) by weight of an anionic synthetic surfactant as previously defined;

(b) from 0 to about 12% (preferably from about 0.5 to about 4%) by weight of a cosurfactant selected from the group consisting of:

(i) quaternary ammonium surfactants having the formula:

30 [R2(OR3),][R4(OR3),]2 R5 N+X-

wherein R², each R³, R⁴, R⁵, X and y are as previously defined; (ii) diguaternary ammonium surfactants having the formula:

35 [R²(OR³)₂][R⁴(OR³)₂]₂N⁺R³N⁺R⁵[R⁴(OR³)₂]₂ (X⁻)₂

wherein R^2 , R^3 , R^4 , y and X are as defined above; particularly preferred are the C_8 - C_{16} alkyl pentamethylethylenediamine chloride, bromide and methylsulfate salts;

(iii) amine surfactants having the formula:

[R2(OR3),][R4(OR3),]R5N

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wherein R², R³, R⁴, R⁵ and y are as defined above; particularly preferred are the C₁₂-C₁₅ alkyl dimethyl amines;

(iv) diamine surfactants having the formula:

[R2(OR3),][R4(OR3),]NR3NR5[R4(OR3),]

wherein R², R³, R⁴, R⁵ and y are as defined above; particularly preferred are the C₁₂-C₁₆ alkyl dimethyl diamines:

(v) amine oxide surfactants having the formula:

[R2(OR3),][R4(OR3),]R5N→O

wherein R², R³, R⁴, R⁵ and y are as defined above; particularly preferred are the C₁₂-C₁₅ alkyldimethyl amine oxides; and (vi) di(amine oxide) surfactants having the formula:

$$[{{\mathsf{R}}^{2}}{{\left({{\mathsf{OR}}^{3}} \right)}_{y}}][{{\mathsf{R}}^{4}}{{\left({{\mathsf{OR}}^{3}} \right)}_{y}}]{{\mathsf{NR}}^{3}}{{\mathsf{NR}}^{5}}[{{\mathsf{R}}^{4}}{{\left({{\mathsf{OR}}^{3}} \right)}_{y}}]$$

wherein R², R³, R⁴, R⁵ and y are as defined above; preferred are the C₁₂-C₁₆ alkyl trimethylethylene di(amine oxides) and

(c) from 0 to about 40% by weight (preferably 5 to about 30% by weight, and most preferably from about 10 to 20% by weight) of a fathy acid containing from about 10 to about 22 carbon atoms (preferably a Ct₁₀-Ct₁₄ saturated fathy acid or mixtures thereof), the mole ratio of the anionic surfactant to the cosultractant being at least 1 and preferably from about 2:1 to about 20:1.

Such compositions also preferably contain from about 3 to about 15% by weight of an ethoxylated alcohol or ethoxylated alkyl phenol (nonionic surfactants) as previously defined. Highly preferred compositions of this type also preferably contain from about 2 to about 10% by weight (preferably from about 3 to about 8% by weight) of a water-soluble polycarboxylate builder (preferably citric acid) and minor amounts (e.g., less than about 20% by weight) of neutralizing agents, buffering agents, phase regulants, hydrotropes, enzymes, enzymes stabilizing agents, polyacids, suds regulants, opacifiers, antioxidants, bactericides, dyes, perfumes and brighteners, such as those described in U.S. Patent 4,285,841 to Barrat et al., Issued August 25, 1981.

B. Detergent Formulations Containing Certain Anionic Surfactants

When high levels of certain anionic detergent surfactants are used, the compounds of the present invention may not deposit as well on the fabric during laundering. See U.S. Patent 4.116.88 to Derstadt et al., Issued September 26, 1978, which describes the incompatibility of ethylene terphthalate/PEG terephthalate soil release polyesters with certain anionic detergent surfactants. These anionic surfactants include the alkyl sulfates and particularly the alkyl benzene sulfonates. Inclusion of certain detergent builders used of uniform the properties of the description of certain detergent builders used to determine the description of the properties. The properties of t

This decreased performance can be offset by inclusion of higher levels of nonionic detergent surfactants, i.e. above about 50% by weight of the surfactant system. However, higher levels of nonionic surfactants do not provide as good cleaning as anionic surfactants, especially in granular detergent formulations. Accord35 ingly, inclusion of a small amount (e.g. from about 0.5 to about 2% by weight of the total composition) of a cationic detergent surfactant(s) as previously described can be used to improve the soil release performance of the compounds. Also, soil release performance can be boosted by simply including more of the compounds of the present invention.

40 Specific Embodiments of Detergent Compositions According to the Present Invention

Embodiment I

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The following embodiments illustrate detergent compositions of the present invention: A granular detergent composition is as follows:

	Component	Wt. % .
	Polyester of Example 1 *	2.0
5	Sodium C ₁₄ -C ₁₅ alkylethoxysulfate	10.7
	C ₁₃ linear alkyl benzene sulfonic acid	4.3
	C ₁₂ -C ₁₄ alkylpolyethoxylate (6)	0.5
10	C ₁₂ alkyltrimethyl ammonium chloride	0.5
10	Sodium toluene sulfonate	1.0
	Sodium tripolyphosphate	32.9
	Sodium carbonate	20.3
15	Sodium silicate	5.8
	Minors and water	Balance to 100

* Enrobed in PEG having an average M.W. 8,000.

Except for the enrobed polyester particles, the components are added together with continuous mixing to form an aqueous slurry which is then spray dried to form granules. The enrobed polyester particles are then mixed with the granules to form the composition.

25 Embodiment II

A liquid detergent composition is formulated as follows:

	Component	Wt. %
30	Polyester of Example 2	1.0
	PEA ₁₈₉ E ₁₇ *	1.0
	Sodium C ₁₂ alkylethoxy (1) sulfate	9.4
35	C ₁₂ -C ₁₃ alcohol polyethoxylate (6.5)	21.5
	Ethanol	7.5
	Sodium diethylenetriamine pentaacetate	0.2
40	MAXATASE	0.026 Anson
		units/g
	TERMAMYL	0.51 KNu/g
	Sodium formate	1.6
45	Calcium formate	0.1
	Minors and water	Balance to 100
		diames of athornilation

 \star Polyethyleneamine having M.W. of 189 and degree of ethoxylation of 17 at each reactive hydrogen.

The components are added together with continuous mixing to form the composition.

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Embodiments III and IV

Liquid detergent compositions are as follows:

5	Component	Wt. 1	
		III	IV
	Polyester of Example 2	1.0	1.0
10	PEA ₁₈₉ E ₁₇	2.0	1.5
	C ₁₄ -C ₁₅ alkylpolyethoxy (2.25) sulfuric acid	12.0	10.8
	C ₁₃ linear alkylbenzene sulfonic acid	8.0	8.0
	C ₁₂ alkyl trimethylammonium chloride	0.6	1.2
15	C ₁₂ -C ₁₃ alcohol polyethoxylate (6.5)	5.0	6.5
	Coconut fatty acid	10.0	13.0
	Oleic acid	-	2.0
20	Citric acid monohydrate	4.0	4.0
	Diethylenetriamine pentaacetic acid	0.2	0.2
	Protease enzyme	0.8	0.8
25	Amylase enzyme	0.2	0.2
	Monoethanolamine	2.0	2.0
	Sodium hydroxide	2.4	1.7
30	Potassium hydroxide	1.1	2.7
30	1,2-Propanediol	3.5	7.3
	Ethanol	8.5	7.8
	Formic acid	0.08	0.7
35	Boric acid	1.3	
	Calcium ion	0.03	0.03
	Minors and water	Balance	to 100

Embodiment III is prepared by adding the components together with continuous mixing, in the following order to produce a clear liquid: a paste premix of alkylbenzene sulfonic acid, a portion of the sodium hydroxide, propylene glycol, and a portion of the tending, is paste premix of alkylopleythoxysulfuric acid, a portion of the sodium hydroxide and a portion of the ethanol; pentaacetic acid; a portion of the alcohol polyethoxylate; a premix of water, triethanolamine, brighteners and the remaining ethanic; citric acid; that pack is the remaining ethanol; potassium hydroxide, citric acid; that pack is formic acid, boric acid and calcium; alkyl trimethylammonium chloride; PEA₁₈₃E₁₇ (50% aqueous solution); polyester of Example 1; adjust to pH 8.0; and balance of components. Embodiment IV can be prepared in a similar manner.

Claims

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 An ester compound having branched hydrophilic capping groups and oligomer or polymer backbone, said compound being characterized in that it has the formula:

 $\begin{array}{l} X \{ (OCH_2CH(Y))_n (OR^5)_m \} \{ (A-R^1-A-R^2)_u (A-R^3-A-R^2)_v - \\ -A-R^4-A \{ (R^5O)_m (CH(Y)CH_2O)_n \} X \end{array}$

wherein the A moieties are selected from

-oc-

or

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moieties:

The R¹ moteties are selected from 1,4-phenylene moieties and combinations thereof with from 0 to 50% of other arylene moieties selected from 1,3-phenylene, 1,2-phenylene, 1,8-napthylene, 1,4-napthylene, 2,2-biphenylene, 4,4-biphenylene or mixtures thereof; the R² moieties are ethylene, or substituted ethylene moieties having C₁-C₁ alkyl or alkoxy substituents; the R³ moieties are substituted c₂-C₁ hydrocarbylene moieties having at least one -SO₃M, -COOM, -O-I(R² O)_m(CH(Y)CH₂O)_m}X or -A² (R²-A-R²-A)²-A)²-R²-A crosslinked to another R³ moiety; the R¹ moieties are R¹ or R² moieties, or a mixture thereof; each R² is C₃-C₄ alkylene, or the moiety -R²-A-R²-, wherein R² is a C₁-C₂ alkylene, alkenylene, arylene or alkarylene moiety; each M is H or a water-soluble cation; the Y substituents of each moiety -I(R² O)_m(CH(Y)CH₂O)_m²- are the ether moiety -CH₂(OCH₂CH₂O_bO-X or mixture of said ether moiety and H; each X is H, C₁-C₄ alkyl

-CR7

wherein R^2 is C_1-C_4 alkyl; m and n are numbers such that the moiety $-(CH(Y)CH_2O)$ - comprises at least 50% by weight of the moiety $+(R^2O_m(CH(Y)CH_2O)_3)$, provided that when R^3 is the moiety $+R^2-A$ - R^2 , m is t; each m is at least t; t and t are numbers such that the sum of u+v is from 3 to 25 when w is 0; w is 0 or at least t; and when w is at least t, u, v and w are numbers such that the sum of u+v+v is from 3 to 25.

- A compound according to Claim 1, wherein each R¹ moiety is a 1.4-phenylene moiety; said R² moieties
 comprise from 20% to 100% ethylene moieties. 1.2-propylene moieties or mixtures thereof; said R³
 moieties are selected from substituted C₂-C₁₂ alkylenes,alkenylenes, arylenes, alkarylenes and mixtures thereof; and the Y substituents are a mixture of said -Ch²₂(OCh²₂-Ch²₂)₀OX ether molety and H.
- A compound according to either of Claims 1 or 2, wherein each of said R³ moieties has one -SO₃M substituent.
- 45 4. A compound according to any of Claims 1-3, wherein v is 0, p is 0, m is 0, and n is from 12 to 119.
 - An ester compound having branched hydrophilic capping groups and oligomer or polymer backbone, said compound being characterized in that it has the formula:

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wherein each R¹ moiety is a 1.4-phenylene moiety; the R² moieties are ethylene moieties, 1,2propylene moieties or mixtures thereof; the R³ moieties are potassium or socilum 5-sulfo-1,3-phenylene moieties or substituted 1,3-phenylene moieties having the substituer

$$\begin{array}{c} \circ \\ \circ \\ -\text{CO} + \left(\mathbb{R}^2 - \text{OC} - \mathbb{R}^4 - \text{CO} \right) \xrightarrow{\bullet} & \left(\text{CHCH}_2 \text{O} \right)_{n_1} \left(\text{CH}_2 \text{CH}_2 \text{O} \right)_{n_2} \xrightarrow{\bullet} \times \\ \downarrow \\ \text{CH}_2 \text{OX} \end{array}$$

- at the 5 position; the R⁴ moieties are R¹ or R³ moieties, or a mixture thereof; each X is ethyl or methyl; each n; is from 1 to 5; the sum of each n; h = n; is from 12 to 45; when w is 0, u + v is from 3 to 10; when w is at least 1, u + v + w is from 3 to 10.
- A compound according to Claim 5, wherein v is 0 and said R² moieties comprise from 80 to 100% of ethylene moieties, 1,2-propylene moieties, or mixtures thereof.
 - 7. A detergent composition characterized in that it comprises:
 - (a) from 1 to 75% by weight of a nonionic, anionic, ampholytic, zwitterionic, or cationic detergent surfactant, or mixtures thereof; and
- (b) a soil release component which comprises from 0.01 to 10% of an ester compound according to any of Claims 1-6.
 - 8. A detergent composition according to Claim 7 which comprises from 0.1 to 5% of component (b).
- 40 9. A detergent composition according to either of Claims 7 or 8, wherein component (a) is a nonionic surfactant, an anionic surfactant or a mixture thereof, which comprises from 10 to 50% of said composition.
 - 10. A detergent composition according to any of Claims 7-9 which further comprises from 3 to 30% of a detergent builder and, optionally, which additionally comprises an optical brightener.

Patentansprüche

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 Esterverbindung mit verzweigten hydrophilen Endgruppen und einem oligomeren oder polymeren Rückgrat, welche Verbindung dadurch gekennzeichnet ist, daß sie die Formel:

$$X_{\{(OCH_2CH(Y))_n(OR^5)_m\}}_{\{(A-R^1-A-R^2)_u(A-R^3-A-R^2)_{v'}\}}$$

-A-R⁴-A $_{\{(R^5O)_m(CH(Y)CH_2O)_n\}}_{X}$

55 besitzt, worin die A-Reste unter



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an

-CO-

Resten ausgewählt sind:

die RI-Reste unter 1,4-Phenylenresten und Kombinationen hievon mit 0 bis 50 % anderer Arylenreste ausgewählt sind, welche unter 1,3-Phenylen, 1,2-Phenylen, 1,8-Naphthylen, 1,4-Naphthylen, 2,2-Biphenylen, 4,4-Biphenylen oder Gemischen hievon ausgewählt sind; die R²-Reste Ethylen oder substituierte Ethylenreste mit C₁-C₄-Alkyl- oder -Alkoxysubstituenten derstellen; die R³-Reste substituierte C₂-C_{1s}-Kohlenwasserstolffreste darstellen, welche mindestens einen Substituenten -SO₃M, -COOM, -O-[(R²O_m-(CHY)CH₂O_b-)₁-X oder

-Af (R²-A-R¹-A₂-f.(R²-O_m(CH(Y)CH₂O_m)-X oder mindestens einen Rest -Af (R²-A-R¹-A)₂-f.(R²-A sufweisen, welcher mit einem weiteren R³ vernetzt ist; die R⁴-Reste R¹- oder R³-Reste oder ein Gemisch hievon sind; jeder Rest R¹ to R²-A R³-Reste oder R³-Reste oder R³-Reste oder Alkylen-, -Alkenylen-, Arylen- oder Alkarylennst darstellt; jeder Rest M H oder ein wasserlösliches Kation bedeutet; die Y-Substituenten jedes Restes -{(R²-O_m(CH(Y)CH₂O_m)-} für den Eiherrest -CH₂-(CCH₂-CH₃-)_C-Y- oder ein Gemisch aus dem genannten Etherrest und H stehen; jeder Rest X H, C₁-C₄-Alkyl oder



bedeutet, worin R⁷ C₁-C₄-Alkyl ist; m und n solche Zahlen sind, daß der Rest -(CH(Y)CH₂O)-mindestens 50 Gew.-% des Restes {R⁶D_{in}(R(Y)CH₂O)_h} unfaßt, mit der Maßgabe, daß, wenn R⁶ den Rest -R⁶ -ArsR⁶ darstellt, m den Wert 1 besitzt; jedes n mindestens 6 ist; p 0 oder mindestens 1 beträgt; u und v solche Zahlen sind, daß die Summe von u + v von 3 bis 25 beträgt, wenn w den Wert 0 besitzt; w 0 oder mindestens 1 ist; und für den Fall, daß w mindestens 1 beträgt, u, v und w solche Zahlen sind, daß die Summe von u + v + w on 3 bis 25 beträgt.

- 2. Verbindung nach Anspruch 1, worin jeder R¹-Rast ein 1,4-Phenylenrest ist; die genannten R²-Reste 20% bis 100% Ethylenreste, 1,2-Propylenreste oder Gemische hievon umfassen; die genannten R³-Reste unter substitutierten C₂-C₁₂-Alkiylenen, -Alkenylenen, Alylenen, Alkenylenen und Gemischen hievon ausgewählt sind; und die Y-Substituenten ein Gemischa aus dem genannten -CH₂(OCH₂CH₂)₀O-X-Etherrestund H sind.
- Verbindung nach einem der Ansprüche 1 oder 2, worin jeder der genannten R²-Reste einen -SO₃ M Substituenten aufweist.
 - Verbindung nach einem der Ansprüche 1 bis 3, worin v 0 ist, p 0 beträgt, m 0 ist und n von 12 bis 119 beträgt.
- Esterverbindung mit verzweigten hydrophilen Endgruppen und einem oligomeren oder polymeren Rückgrat, welche Verbindung dadurch gekennzeichnet ist, daß sie die Formel:

besitzt, worin jeder R¹-Rest ein 1,4-Phenylenrest ist, die R²-Reste Ethylenreste, 1,2-Propylenreste oder Gemische hievon darstellen; die R³-Reste Kalium- oder Natrium-5-sulfo-1,3-phenylenreste oder substituierte 1,3-Phenylenreste darstellen, welche an der 5-Stellung den Substituenten

$$-\frac{0}{\text{co}_{-1-}} (R^2 - \text{oc}_{-R}^4 - \frac{0}{\text{co}}) - \frac{1}{1 - \frac{1}{1}} (\text{chch}_{2}\text{o})_{n_1} (\text{ch}_{2}\text{ch}_{2}\text{o})_{n_2} - x$$

aufweisen; die R⁴-Reste R¹- oder R³-Reste oder ein Gemisch hievon sind; jeder Rest X Ethyl oder Methyl bedeutet; jedes n; von 1 bis 5 beträgt; die Summe von jedem n; +n; von 12 bis 45 beträgt; wenn v 0 ist, u+v von 3 bis 10 beträgt; m Fall, daß w mindestens 1 ist, u+v +v von 3 bis 10 beträgt;

- Verbindung nach Anspruch 5, worin v 0 ist und die genannten R²-Reste 80 bis 100% an Ethylenresten, 1,2-Propylenresten oder Gemischen hievon umfassen.
- Detergenezusammensetzung, dadurch gekennzeichnet, daß eie:

 (a) 1 Gew.-% bis 75 Gew.-% eines nichtionischen, anionischen, ampholytischen, zwitterionischen oder kationischen grenzflächensatiliven Detergensmittels oder von Gemischen hievori; und
 (b) eine Schmutzlösekomponente umfaßt, welche 0,01% bis 10% einer Esterverbindung nach einem der Ansprüche 1 bis 6 enthält.
 - 8. Detergenszusammensetzung nach Anspruch 7, welche 0,1 bis 5% der Komponente (b) umfaßt.
 - Detergenszusammensetzung nach einem der Ansprüche 7 oder 8, worin die Komponente (a) ein nichtlichisches grenzflächenaktives Mittle, ein anionisches grenzflächenaktives Mittel oder ein Gemisch hievon ist, welches 10% bis 50% der genannten Zusammensetzung darstellt.
- 45 10. Detergenszusammensetzung nach einem der Ansprüche 7 bis 9, welche ferner 3% bis 30% eines Detergensgerüststoffes und wahlweise zusätzlich einen optischen Aufheller umfaßt.

Revendications

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50 1. Composé ester possédant des groupes bloquants ramifiés hydrophiles et une chaîne principale oligomère ou polymère, ledit composé étant caractérisé en ce qu'il répond à la formule :

$$X[(OCH_2CH(Y))_n(OR^5)_m] + (A-R^1-A-R^2)_u(A-R^3-A-R^2)_{v'}]$$

-A-R⁴-A $[(R^5O)_m(CH(Y)CH_2O)_n] \times$

dans laquelle les groupements A sont choisis parmi les groupements



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les groupements R¹ sont choisis parmi les groupements 1,4-phénylène et leurs combinaisons avec de 0 à 50% d'autres groupements anylène choisis parmi le 1,3-phénylène, le 1,2-phénylène, le 1,2-phénylène, le 4,4-biphénylène, le 4,4-biphénylène, le 4,4-biphénylène, le 4,5-phénylène, le 4,5-phénylène, le 4,5-phénylène, le 4,5-phénylène, le 4,5-phénylène, le 4,5-phénylène, le 1,8-phénylène, l

-A.[(R²-A-R¹-A)],-[(R²O)],(CH(Y)CH₂O),]-X ou au moins un groupement :A.[(R²-A-R¹-A)],-R²-A réticulé à un autre groupement R²; les groupements R¹ sont des groupements R¹ ou R², ou un mélange de ceux-cl; chaque R² est un groupement alkylène, en C₂-C₂, ou le groupement -R²-A-R², dans loquel R² est un groupement alkylène, alcénylène, arylène ou alkylarylène en C₁-C₁; chaque M est H ou un cation hydrosoluble; les substituants Y de chaque groupement (†R²O)_m(CH(Y)CH₂O)_m-S ont le groupement éther -CH₂(OCH₂CH₂)_pO-X ou un mélange dudit groupement éther et de H; chaque X est H, un groupe alkyle on C₁-C₄ ou

dans lequel R' est un groupe alkyle en $C_1 \cdot C_4$; m et n sont des nombres tels que le groupement $-(SH^*)$ ($C_1 \cdot C_2 \cdot C_3 \cdot C_4 \cdot C_4$

- 2. Composé selon la revendication 1, dans lequel chaque groupement R¹ est un groupement 1.4-phényihen; lesdits groupements R² comprennent de 20% à 100% de groupements éthyihen, de groupements 11.2-propylène, ou de mélanges de ceux-ci; lesdits groupements P¹ sont choisis parmi des groupes alkylène, alcényiène, arylène ou alkylarylène substitués en Cz-O₁z, et des mélanges de ceux-ci; et les substituants Y sont un mélange dudit groupement éther -CH₂(OCH, Eh₂)b-X et de 1.
- Composé selon l'une quelconque des revendications 1 ou 2, dans lequel chacun desdits groupements R³ possède un substituant -SO₃M.
 - Composé selon l'une quelconque des revendications 1-3, dans lequel v est nul, p est nul, m est nul et n a une valeur de 12 à 119.
 - 5. Composé ester possédant des groupes bloquants ramifiés hydrophiles et une chaîne principale oligomère ou polymère, ledit composé étant caractérisé en ce qu'il répond à la formule :

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dans laquelle chaque groupement R¹ est un groupement 1,4-phénylène; les groupements R² sont des groupements éthyène, des groupements 1,2-propylène ou des mélanges de ceux-ci; les groupements R³ sont des groupements 5-sulfo-1,3-phénylène potassique ou sodique ou des groupements 1,3phénylène substitués possédant le substituant

$$\begin{array}{c} \bigcap\limits_{-\text{CO}-\text{I}-}^{\bigcap} (\mathbb{R}^2 - \mathbb{O}\mathbb{C} - \mathbb{R}^4 - \mathbb{O}\mathbb{O}) \xrightarrow{\text{I} \quad \text{w. f}} (\text{CHCH}_2\text{O})_{n_1} (\text{CH}_2\text{CH}_2\text{O})_{n_2} \xrightarrow{\text{I} \quad \text{CH}_2\text{OX}} \\ \bigcap\limits_{\text{CH}_2\text{OX}} (\mathbb{C} + \mathbb{C} + \mathbb{C$$

en position 5; les groupements R⁴ sont des groupements R¹ ou R³, ou un mélange de ceux-ci; chaque X est un groupe éthyle ou méthyle; chaque n. a une valeur de 1 à 5; la somme n₁ + n₂ a une valeur de 12 à 45; lorsque w est nul, u + v a une valeur de 3 à 10; lorsque w est au moins égal à 1, u + v + w a une valeur de 3 à 10.

- Composé selon la revendication 5, dans lequel v est nul et lesdits groupements R² comprennent de 80
 à 100% de groupements éthylène, de groupements 1,2-propylène, ou de mélanges de ceux-ci.
 - 7. Composition détergente, caractérisée en ce qu'elle comprend :
 - (a) de 1 à 75% en poids d'un tensioactif détergent non ionique, anionique, ampholyte, zwittérionique ou cationique, ou d'un mélance de ceux-ci: et
 - (b) un constituant de libération des salissures qui comprend de 0,01 à 10% d'un composé ester selon l'une quelconque des revendications 1-6.
 - 8. Composition détergente selon la revendication 7, qui comprend de 0,1 à 5% de constituant (b).
- 9. Composition détergente selon l'une quelconque des revendications 7 ou 8, dans laquelle le constituant (a) est un tensioactif non ionique, un tensioactif anionique, ou un mélange de ceux-ci, qui constitue de 10 à 50% de ladite composition.
- 10. Composition détergente selon l'une quelconque des revendications 7-9, qui comprend aussi de 3 à 30% d'un adjuvant de détergence et, éventuellement, qui comprend aussi un azurant optique.

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